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PASSWORD:

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* * *	* *	* *	* *	* Welcome to STN International * * * * * * * * *
				Welcome to SIN International
NEWS	1			Web Page for STN Seminar Schedule - N. America
NEWS	2	AUG	06	CAS REGISTRY enhanced with new experimental property tags
NEWS	3	AUG	06	FSTA enhanced with new thesaurus edition
NEWS	4	AUG	13	CA/CAplus enhanced with additional kind codes for granted
				patents
NEWS	5	AUG	20	CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS	6	AUG	27	Full-text patent databases enhanced with predefined
				patent family display formats from INPADOCDB
NEWS		AUG		USPATOLD now available on STN
NEWS	8	AUG	28	CAS REGISTRY enhanced with additional experimental
	_			spectral property data
NEWS	9	SEP	0.7	STN AnaVist, Version 2.0, now available with Derwent
NEWS	2.0	SEP	1.0	World Patents Index FORIS renamed to SOFIS
NEWS		SEP		INPADOCDB enhanced with monthly SDI frequency
NEWS		SEP		CA/CAplus enhanced with monthly SDI frequency
MEMP	12	JEE	1,	1967-1998
NEWS	13	SEP	17	CAplus coverage extended to include traditional medicine
				patents
NEWS		SEP		EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS	15	OCT	02	CA/CAplus enhanced with pre-1907 records from Chemisches
NEWS	16	OCT	10	Zentralblatt BEILSTEIN updated with new compounds
NEWS		NOA		Derwent Indian patent publication number format enhanced
NEWS		NOV		WPIX enhanced with XML display format
NEWS		NOV		ICSD reloaded with enhancements
NEWS			04	LINPADOCDB now available on STN
NEWS			14	BEILSTEIN pricing structure to change
NEWS	22	DEC	17	USPATOLD added to additional database clusters
NEWS	23	DEC	17	IMSDRUGCONF removed from database clusters and STN
NEWS	24	DEC	17	DGENE now includes more than 10 million sequences
NEWS	25	DEC	17	TOXCENTER enhanced with 2008 MeSH vocabulary in
				MEDLINE segment
NEWS		DEC		MEDLINE and LMEDLINE updated with 2008 MeSH vocabulary
NEWS		DEC		CA/CAplus enhanced with new custom IPC display formats
NEWS	28	DEC	17	STN Viewer enhanced with full-text patent content
NELLO	00	77.11	0.0	from USPATOLD
NEWS NEWS		JAN JAN		STN pricing information for 2008 now available
MEMP	30	UMIN	10	CAS patent coverage enhanced to include exemplified prophetic substances
NEWS	31	JAN	28	USPATFULL, USPAT2, and USPATOLD enhanced with new
				custom IPC display formats
NEWS		JAN		MARPAT searching enhanced
NEWS	33	JAN	28	USGENE now provides USPTO sequence data within 3 days
NEWS	3.4	JAN	20	of publication TOXCENTER enhanced with reloaded MEDLINE segment
MEMP	34	OMIN	40	TONGERIER EIMENCEG WITH TETOGGEG MEDEINE SEGMENT

NEWS 35 JAN 28 MEDLINE and LMEDLINE reloaded with enhancements NEWS 36 FEB 08 STN Express, Version 8.3, now available

NEWS EXPRESS FEBRUARY 08 CURRENT WINDOWS VERSION IS V8.3, AND CURRENT DISCOVER FILE IS DATED 24 JANUARY 2008

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FILE 'HOME' ENTERED AT 11:40:46 ON 13 FEB 2008

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ENTRY SESSION
FULL ESTIMATED COST 0.63
0.63

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http://www.cas.org/support/stngen/stndoc/properties.html

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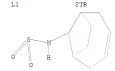


chain nodes:
10 11 12 13 14
ring nodes:
1 2 3 4 5 6 7 8 9
chain bonds:
3-10 10-11 10-12 12-13 12-14
ring bonds:
1-2 1-7 2-3 2-9 3-4 4-5 4-8 5-6 6-7 8-9
exact/norm bonds:
2-9 3-10 4-8 10-12 12-13 12-14
exact bonds:
1-2 1-7 2-3 3-4 4-5 5-6 6-7 8-9 10-11
isolated ring systems:
containing 1:

Match level: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 11:CLASS 13:CLASS 14:CLASS

L1 STRUCTURE UPLOADED

=> d l1 L1 HAS NO ANSWERS



Structure attributes must be viewed using STN Express query preparation.

=> s 11 SAMPLE SEARCH INITIATED 11:43:41 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 30 TO ITERATE

100.0% PROCESSED 30 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.01

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FULL SEARCH INITIATED 11:43:46 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 479 TO ITERATE

100.0% PROCESSED 479 ITERATIONS

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SEARCH TIME: 00.00.01

13 SEA SSS FUL L1

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SINCE FILE TOTAL ENTRY SESSION COST IN U.S. DOLLARS FILL ESTIMATED COST 178.82 179.45

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L4 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:141037 CAPLUS

DOCUMENT NUMBER: 142:240436

TITLE: Preparation of spirobicyclononenethiadiazole dioxides

and related compounds as γ-secretase inhibitors
INVENTOR(S): Bettati, Michela; Boase, Amanda Louise; Churcher, Ian;

Ladduwahetty, Tamara; Merchant, Kevin John; Quddus, Abdul

PATENT ASSIGNEE(S): Merck Sharp & Dohme Limited, UK

SOURCE: PCT Int. Appl., 39 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PA | TENT | NO. | | | KIN | DATE | | | | | | DATE | | | | | | | | |
|--------|------------------------------|-------|-----|-----|------|-------------|---------|--------|------|------------------------|----------------|------|-------|------|-----|------|-----|--|--|--|
| WC | 0 2005014553 | | | | | A1 20050217 | | | | | WO 2004-GB3277 | | | | | | | | | |
| | W: AE, AG, AL, | | | | AM, | AT, | AU, AZ, | | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH, | | | |
| | | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, | | | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | | | |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, | | | |
| | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, | | | |
| | | TJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UZ, | VC, | VN, | YU, | ZA, | ZM, | ZW | | | |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | | | |
| | | ΑZ, | BY, | KG, | KΖ, | MD, | RU, | ΤJ, | TM, | ΑT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | | | |
| | | EE, | ES, | FI, | FR, | GB, | GR, | HU, | IE, | IT, | LU, | MC, | NL, | PL, | PT, | RO, | SE, | | | |
| | | SI, | SK, | TR, | BF, | ΒJ, | CF, | CG, | CI, | CM, | GA, | GN, | GQ, | GW, | ML, | MR, | NE, | | | |
| | | SN, | TD, | TG | | | | | | | | | | | | | | | | |
| | AU 2004263353 | | | | | | | | | | | | | | | | | | | |
| | 2534 | | | | | | | | | | | | | | | | | | | |
| | 1658 | | | | | | | | | EP 2 | 004- | | 2 | 0040 | 729 | | | | | |
| EF | 1658 | 272 | | | В1 | | 2007 | 0725 | | | | | | | | | | | | |
| | R: | ΑT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IT, | LI, | LU, | NL, | SE, | MC, | PT, | | | |
| | | | | | | | | | | | CZ, | | | | | | | | | |
| CI/ | 1832 | 927 | | | A | | 2006 | 0913 | | CN 2 | 004- | | 2 | 0040 | 729 | | | | | |
| JF | 1832
2007
3680
2289 | 5012 | 06 | | T | | 2007 | 0125 | | JP 2 | 006- | 5223 | 90 | | 2 | 0040 | 729 | | | |
| AT | 3680 | 31 | | | T | | 2007 | 0815 | - 2 | AT 2 | 004- | 7436 | 04 | | 2 | 0040 | 729 | | | |
| ES | 2289 | 537 | | | Т3 | | 2008 | 0201 | 1 | ES 2 | 004- | 7436 | 04 | | 2 | 0040 | 729 | | | |
| | 2006 | | | | | | | | | | | | | | | | | | | |
| | 2006 | | | | A1 | | 2006 | 0824 | | | | | | | | | | | | |
| RIORIT | IORITY APPLN. INFO.: | | | | | | | | | | 003- | | | | | | | | | |
| | | | | | | | | | | | 004-0 | | | | | 0040 | 729 | | | |
| THER S | OHRCE | (8) . | | | CASI | REAC | T 14 | 2 • 24 | 0436 | MA | RPAT | 142 | · 240 | 136 | | | | | | |

OTHER SOURCE(S): CASREACT 142:240436; MARPAT 142:240436

GI

- AB Title compds. [I; n = 0, 1; X = atoms to form a 5-6 membered heteroarom. ring; R5 = (halo-substituted) hydrocarbyl; Ar = (substituted) Ph, 6-membered heteroaryl; Y = bond, NR3; R1 = H; R1R3 = CH2; R2 = (halo-substituted) hydrocarbyl, (substituted) 5-6 membered heteroaryl; R2R3 = atoms to form a (substituted) heterocyclic ring of ≤6 members; R3 = H, alkyl; R4 = halo, alkyl], were prepared as γ-secretase inhibitors (no data). Thus title compound (II) was prepared in several steps from bicyclo[4,2.1]non-3-en-9-one, tert-Bu sulfinamide, F3CCH2NH2, POCl3/DMF, and [5-(4-fluorophenyl)-1-methyl-1H-pyrazol-3-γl]methyltriphenylphosphonium chloride.
- IT 844880-01-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of spirobicyclononenethiadiazole dioxides and related compds. as γ -secretase inhibitors)

II

- RN 844880-01-9 CAPLUS
- CN Sulfamide, N-[3-chloro-4-[(1E)-2-[5-(4-fluorophenyl)-1-methyl-1H-pyrazol-3-yl]ethenyl]bicyclo[4.2.1]non-3-en-9-yl]-N'-propyl- (CA INDEX NAME)

Double bond geometry as shown.

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L4 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2003:133023 CAPLUS

DOCUMENT NUMBER: 138:169963

TITLE: Synthesis of sulfonamido-substituted bridged

bicycloalkyl derivatives for control of beta-amyloid

production

Hannam, Joanne Claire; Harrison, Timothy; Madin,

Andrew; Sparey, Timothy Jason

PATENT ASSIGNEE(S): Merck Sharp & Dohme Limited, UK

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | TENT | | | | KIND | | DATE | | | APP: | LICAT | | DATE | | | | | | |
|---------|------------------------|------|-----|-----|------|-----|------|------|-----|----------------|-------|------|------|----------|-----|----------|-----|--|--|
| | | | | | A1 | _ | | | | WO 2002-GB3559 | | | | | | 20020731 | | | |
| | W: AE, AG, AL, | | | AM, | AT, | AU, | AZ, | BA, | BB | , BG, | BR, | BY, | BZ, | CA, | CH, | CN, | | | |
| | | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC | , EE, | ES, | FI, | GB, | GD, | GE, | GH, | | |
| | | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE | , KG, | KR, | KZ, | LC, | LK, | LR, | LS, | | |
| | | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW | , MX, | MZ, | NO, | NZ, | OM, | PH, | PL, | | |
| | | PT, | RO, | RU, | SD, | SE, | SG, | SI, | SK, | SL | , TJ, | TM, | TN, | TR, | TT, | TZ, | UA, | | |
| | | UG, | US, | UZ, | VN, | YU, | ZA, | ZM, | ZW | | | | | | | | | | |
| | RW: | GH, | GM, | KE, | LS, | MW, | MZ, | SD, | SL, | SZ | , TZ, | UG, | ZM, | ZW, | AT, | BE, | BG, | | |
| | | CH, | CY, | CZ, | DE, | DK, | EE, | ES, | FI, | FR | , GB, | GR, | IE, | IT, | LU, | MC, | NL, | | |
| | | PT. | SE. | SK, | TR. | BF, | BJ, | CF. | CG, | CI | CM, | GA, | GN, | GO, | GW, | ML. | MR. | | |
| | | NE. | SN, | TD, | TG | | | | | | | | | | | | | | |
| AU | 2002 | 3553 | 59 | | | | | | | AU : | 2002- | 3553 | | 20020731 | | | | | |
| US | 2004 | 1861 | 47 | | A1 | | 2004 | 0923 | | US 2004-484290 | | | | | | 20040120 | | | |
| US | 7205 | 434 | | | B2 | | 2007 | 0417 | | | | | | | | | | | |
| PRIORIT | PRIORITY APPLN. INFO.: | | | | | | | | | GB : | 2001- | 1915 | 2 | | A 2 | 0010 | 806 | | |
| | | | | | | | | | | WO : | 2002- | GB35 | 59 | | W 2 | 0020 | 731 | | |
| OTHER S | | | | | | | 138: | 1699 | 63 | | | | | | | | | | |

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Title compds. I [A,B = together with the carbon atoms bonded to L1R4 and H complete a (un)substituted ring containing 5-10 carbon atoms; R1 = H, alkyl, alkenyl; R2 = H, acyl; R3 = alkyl, cycloalkyl, alkenyl, alkynyl, aryl, etc.; R4 = H, halo, aryl, heterocyclyl, CN, alkoxy, amino, etc.; L1 = bond, alkylene, etc.] are prepared For instance, Et cyclopentanone-2carboxylate was reacted with o-xylylene dibromide (DMF, NaOEt) and the resulting adduct treated with LDA in THF at -78° to give II. II was treated in the following manner: i. THF, H2NOH+HCl, NaOAc; ii. HOAc, H2-PtO; iii. CH2Cl3, Et3N, 5-chlorothiophenesulfonyl chloride and iv. THF, LAH to provide sulfonamide III. I modulate the production of B-amyloid from amyloid precursor protein and are useful in the treatment of Alzheimer's disease.

497862-61-0P

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(synthesis of sulfonamido-substituted bridged bicycloalkyl derivs. for control of beta-amyloid production)

RN 497862-61-0 CAPLUS

CN Bicyclo[4.2.1]non-3-ene-1-carboxylic acid, 9-[[(5-chloro-2thienyl)sulfonyl]amino]-, ethyl ester, (1R,6R,9S)-rel- (CA INDEX NAME)

IT 497862-62-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(synthesis of sulfonamido-substituted bridged bicycloalkyl derivs. for control of beta-amyloid production)

RN 497862-62-1 CAPLUS

CN 2-Thiophenesulfonamide, 5-chloro-N-[(1R,6R,9S)-1-(hydroxymethyl)bicyclo[4.2.1]non-3-en-9-yl]-, rel- (CA INDEX NAME)

REFERENCE COUNT:

4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:353420 CAPLUS DOCUMENT NUMBER: 136:369505

TITLE: Synthesis of sulfonamido-substituted bridged

bicycloalkyl derivatives as γ-secretase

inhibitors

INVENTOR(S): Collins, Ian James; Hannam, Joanne Claire; Harrison, Timothy; Lewis, Stephen John; Madin, Andrew; Sparey,

Timothy Jason; Williams, Brian John

PATENT ASSIGNEE(S): Merck Sharp & Dohme Limited, UK

PCT Int. Appl., 151 pp.

SOURCE:

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

| | | | | | | | | | APPLICATION NO. | | | | | | | | | | |
|-------|----------------|------|------|------|-----|----------------|-----|------|-----------------|----------------|-------|----------------|------|-----|------|------|------|-----|--|
| | | | | | | | | | | WO 2001-GB4817 | | | | | | | | | |
| | W: AE, AG, AL, | | | AL, | AM, | AT, | AU, | AZ, | BA, | BB | , BG, | BR, | BY, | ΒZ, | CA, | CH, | CN, | | |
| | | | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC | , EE, | ES, | FI, | GB, | GD, | GE, | GH, | |
| | | | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE | , KG, | KR, | KZ, | LC, | LK, | LR, | LS, | |
| | | | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW | , MX, | MZ, | NO, | NZ, | PH, | PL, | PT, | |
| | | | RO, | RU, | SD, | SE, | SG, | SI, | SK, | SL, | TJ | , TM, | TR, | TT, | TZ, | UA, | UG, | US, | |
| | | | UZ, | VN, | YU, | ZA, | ZW | | | | | | | | | | | | |
| | | RW: | GH, | GM, | KE, | LS, | MW, | MZ, | SD, | SL, | SZ | , TZ, | UG, | ZW, | ΑT, | BE, | CH, | CY, | |
| | | | DE, | DK, | ES, | FI, | FR, | GB, | GR, | IE, | ΙT | , LU, | MC, | NL, | PT, | SE, | TR, | BF, | |
| | | | | | | | | | | | | , ML, | | | | | | | |
| | CA | 2427 | 206 | | | A1 | | 2002 | 0510 | | CA | 2001- | | 2 | 0011 | 029 | | | |
| | ΑU | 2002 | 0107 | 47 | | A | | 2002 | 0515 | AU 2002-10747 | | | | | | | | | |
| | | | | | | EP 2001-978652 | | | | | | | 0011 | 029 | | | | | |
| | EΡ | 1334 | 085 | | | B1 | | 2005 | 0824 | | | | | | | | | | |
| | | R: | | | | | | | | | | , IT, | | LU, | NL, | SE, | MC, | PT, | |
| | | | | | | | | | | | | , TR | | | | | | | |
| | JΡ | 2004 | 5131 | 8 0 | | T | | 2004 | 0430 | | JP | 2002- | 5393 | 15 | | 2 | 0011 | 029 | |
| | JΡ | 3880 | 051 | | | B2 | | 2007 | 0214 | | | 2001-
2001- | | | | | | | |
| | ΑT | 3027 | 53 | | | T | | 2005 | 0915 | | ΑT | 2001- | 9786 | 52 | | 2 | 0011 | 029 | |
| | ES | 2248 | 397 | | | T3 | | 2006 | 0316 | | ES | 2001- | 9786 | 52 | | 2 | 0011 | 029 | |
| | US | 2004 | 0490 | 38 | | A1 | | 2004 | 0311 | US 2003-415751 | | | | | | 2 | 0030 | 501 | |
| | US | 7138 | 400 | | | B2 | | 2006 | 1121 | | | 2006- | | | | | | | |
| | | | | | | A | | 2006 | 0914 | | JP | 2006- | 7813 | 6 | | 2 | 0060 | 322 | |
| PRIOR | IT | APP | LN. | INFO | .: | | | | | | GB | 2000- | 2682 | 7 | | A 2 | 0001 | 102 | |
| | | | | | | | | | | | GB | 2001- | 2268 | 5 | | A 2 | 0010 | 920 | |
| | | | | | | | | | | | JP | 2002- | 5393 | 15 | | A3 2 | 0011 | 029 | |
| | | | | | | | | | | | WO | 2001- | GB48 | 17 | | W 2 | 0011 | 029 | |

OTHER SOURCE(S): MARPAT 136:369505

GI

AB Title compds. I [A, B = (CXY)p, (CXY)qCY=CY(CXY)r, (CXY)xNR13(CXY)y, etc.; X = halo, R9, OR9, SR9, S(O)1-2R10, OSO2R9, N(R9)2, COR9, CO2R9, etc.; Y = H, alkyl or X, Y together = O, S, N-OR11, CHR11; provided neither A nor B comprises more than one CXY moiety which is other than CH2; p = 1-6; q, r, x, v = 0-2; provided that at least one of A and B comprises a chain of 2 or more atoms, such that the ring completed by A and B contains at least 5 atoms; R1 = H, alk(en)yl or R1 and R15 together may complete a 5-, 6- or 7-membered cyclic sulfamide; R2 = H, C1, alkyl, aryl, aryl-alkyl, cycloalkyl, acyl, etc.; R9 = H or R10 or two R9 groups together with a nitrogen atom to which they are mutually attached may complete a pyrrolidine, piperidine, piperazine, etc.; R10 = alkyl, perfluoroalkyl, cycloalkyl, etc.; R11 = H, alkyl, etc.; R14 = H, alkyl, etc.; R15 = H, alkyl or R15 and R1 together complete a 5-, 6- or 7-membered cyclic sulfamide] were prepared Over 150 synthetic examples were disclosed. For instance, prior art amine II was sulfonylated with catechol sulfate and the intermediate treated with n-PrNH2 (dioxane, 80°C, 1 h) to give III. I are inhibitors of γ -secretase and are cytotoxic with EC50 < 10 µM for human app695. Compds. of the invention are useful in the treatment of and/or prevention of Alzheimer's disease. 423167-24-2P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug; synthesis of sulfonamido-substituted bridged bicycloalkyl derivs. as γ -secretase inhibitors)

RN 423167-24-2 CAPLUS

CN Sulfamide, N-(9-syn)-bicyclo[4.2.1]non-3-en-9-y1-N'-propyl- (CA INDEX NAME)

Relative stereochemistry.



IT 423168-72-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; synthesis of sulfonamido-substituted bridged bicycloalkyl derivs. as γ-secretase inhibitors)

RN 423168-72-3 CAPLUS

CN Sulfamic acid, [(9-syn)-bicyclo[4.2.1]non-3-en-9-yl]-, 2-hydroxyphenyl ester (9CI) (CA INDEX NAME)

Relative stereochemistry.



REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:713298 CAPLUS

DOCUMENT NUMBER: 135:272746

TITLE: Synthesis of sulfonamido-substituted bridged

bicycloalkyl derivatives as γ-secretase

inhibitors

INVENTOR(S): Belanger, Patrice Charles; Collins, Ian James; Hannam, Joanne Claire; Harrison, Timothy; Lewis, Stephen John;

Joanne Claire; Harrison, Timothy; Lewis, Stephen John; Madin, Andrew; McIver, Edward Giles; Nadin, Alan John; Neduvelil, Joseph George; Shearman, Mark Steven;

JP 2002-539315 A3 20011029

Smith, Adrian Leonard; Sparey, Timothy Jason; Stevenson, Graeme Irvine; Teall, Martin Richard

PATENT ASSIGNEE(S): Merck Sharp & Dohme Limited, UK; Merck Frosst Canada +

SOURCE: PCT Int. Appl., 199 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| | | | | | | KIND DATE | | | | | LICAT | | | | | | | |
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| | WO 2001070677 | | | | | | | | | | | | | | | | | |
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| | | | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG | , KP, | KR, | KZ, | LC, | LK, | LR, | LS, |
| | | | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW | , MX, | MZ, | NO, | NZ, | PL, | PT, | RO, |
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| | | | VN, | YU, | ZA, | ZW | | | | | | | | | | | | |
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| | | | | | | | | | | | | , MR, | | | | | | |
| | CA 2404125 | | | | | | | | | | | | | | | | | |
| | EP 1268412 | | | | | | | | E | EΡ | 2001- | 9119 | | 2 | 0010 | 315 | | |
| Ε | P | 1268 | 412 | | | B1 | 2006 | 1122 | | | | | | | | | | |
| | | R: | | | | | | | | | | , IT, | LI, | LU, | NL, | SE, | MC, | PT, |
| | | | | | | | | | | | | , TR | | | | | | |
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| A | Υ | 3460 | 39 | | | T | | 2006 | 1215 | I | AΤ | 2001- | 9119 | 40 | | 2 | 0010 | 315 |
| Ε | S | 2275 | 657 | | | Т3 | | 2007 | 0616 | E | ES | 2001- | 9119 | 40 | | 2 | 0010 | 315 |
| U | JS | 2004 | 02986 | 62 | | A1 | | 2004 | 0212 | Ţ | US | 2003- | 2392 | 33 | | 2 | 0030 | 205 |
| J | ſΡ | 2006 | 24116 | 63 | | A | | 2006 | 0914 | | JΡ | 2006- | 7813 | 6 | | 2 | 0060 | 322 |
| PRIORI | TY | APP: | LN. : | INFO | . : | | | | | (| GΒ | 2000- | 6717 | | - 2 | A 2 | 0000 | 320 |
| | | | | | | | | | | (| GB 2000-26827 | | | | | A 2 | 0001 | 102 |
| | | | | | | | | | | V | WO | 2001- | GB11. | 54 | 1 | 7 2 | 0010 | 315 |
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OTHER SOURCE(S): MARPAT 135:272746

GI

$$\begin{array}{c} \text{Me} & \overset{\circ}{\longrightarrow} & \overset$$

AB Title compds. I [A, B = (CXY)p; (CXY)qCY:CY(CXY)r; (CXY)xNR13(CXY)y; etc.; X = halo, alkoxy, sulf(a/i/o)nyl, amino, acyl, etc.; Y = H, alkyl; or Xand Y together represent :O, :S, :N-OR, :CH; provided neither A nor B comprises more than one -CXY-moiety which is other than CH; Z completes a (non)aromatic ring system of 5 to 10 atoms, of which 0 to 3 are selected from N, O and S and the remainder are C; Z1 completes a nonarom. ring system of 5 to 10 atoms, of which 0 to 3 are independently selected from O, N and S and the remainder are C; Z2 completes a 5- or 6-membered heteroaryl ring; m, n = 0 - 1; p = 1 - 6; q, r, s = 0 - 2; x, y = 0 - 2; provided that when m= n = 0, at least one of A and B comprises a chain of 2 or more atoms, such that the ring completed by A and B contains at least 5 atoms; R1 = H. alkyl, alkenyl; R2 = H, alkyl, aryl(alkyl), cycloalkyl, acyl; R3 = (cyclo)alkyl, alkenyl, alkynyl, (hetero)arylalkyl, etc.] were prepared Over 270 synthetic examples were disclosed. For instance, 1,2-Bis(bromomethyl)benzene was added to 1-cyclopent-1-enylpyrrolidine (CH3CN, (i-Pr)2NEt) to give iminum bromide II. II was converted to the oxime (EtOHag, NH2OH, NaOAc); the oxime was reduced (HOAc, PtO2, H2 @ 30 psi, 2 h) and the resulting amine sulfonylated (DCM, pyridine, p-TsCl, 16 h) to give III. I are inhibitors of γ -secretase and are cytotoxic with EC50 < 10 µM for human app695. Compds. of the invention are useful in the treatment of and/or prevention of Alzheimer's disease. 362654-13-5P 362654-14-6P 362654-15-7P 362654-16-8P 362654-17-9P 362654-66-8P 362654-67-9P 362654-68-0P RL: BAC (Biological activity or effector, except adverse); BSU (Biological

III

RN 362654-13-5 CAPLUS CN Benzenesulfonamide, N-(9-syn)-bicyclo[4.2.1]non-3-en-9-yl-4-methyl- (CA INDEX NAME)

BIOL (Biological study); PREP (Preparation); USES (Uses)

derivs. as y-secretase inhibitors)

study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);

(drug; synthesis of sulfonamido-substituted bridged bicycloalkyl

Relative stereochemistry.

RN 362654-14-6 CAPLUS

CN Benzenesulfonamide, N-(9-syn)-bicyclo[4.2.1]non-3-en-9-yl-4-fluoro- (CA INDEX NAME)

Relative stereochemistry.

RN 362654-15-7 CAPLUS

CN Benzenesulfonamide, N-(9-syn)-bicyclo[4.2.1]non-3-en-9-yl- (CA INDEX NAME)

Relative stereochemistry.

RN 362654-16-8 CAPLUS

CN 2-Thiophenesulfonamide, N-(9-syn)-bicyclo[4.2.1]non-3-en-9-yl- (CA INDEX NAME)

Relative stereochemistry.

RN 362654-17-9 CAPLUS

CN 2-Thiophenesulfonamide, N-(9-syn)-bicyclo[4.2.1]non-3-en-9-yl-5-chloro-(CA INDEX NAME)

Relative stereochemistry.

RN 362654-66-8 CAPLUS

CN Benzenesulfonamide, N-[(9-syn)-9-methylbicyclo[4.2.1]non-3-en-9-yl]- (CA INDEX NAME)

Relative stereochemistry.

RN 362654-67-9 CAPLUS

CN 2-Thiophenesulfonamide, 5-chloro-N-[(9-syn)-9-methylbicyclo[4.2.1]non-3-en-9-yl]- (CA INDEX NAME)

Relative stereochemistry.

RN 362654-68-0 CAPLUS

CN 2-Thiophenesulfonamide, 5-chloro-N-[(9-syn)-9-(2propenyl)bicyclo[4.2.1]non-3-en-9-y1]- (9CI) (CA INDEX NAME)

Relative stereochemistry.

REFERENCE COUNT:

13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

(FILE 'HOME' ENTERED AT 11:40:46 ON 13 FEB 2008)

FILE 'REGISTRY' ENTERED AT 11:42:36 ON 13 FEB 2008

STRUCTURE UPLOADED

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